Southern Implants (Pty) Ltd
% Kevin Walls
Consultant
Regulatory Insight, Inc.
33 Golden Eagle Lane
Littleton, Colorado 80127

Re: K172160
  Trade/Device Name: Southern Implants PEEK Abutments
  Regulation Number: 21 CFR 872.3630
  Regulation Name: Endosseous Dental Implant Abutment
  Regulatory Class: Class II
  Product Code: NHA
  Dated: January 5, 2018
  Received: January 8, 2018

Dear Kevin Walls:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820);
and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (https://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/) and CDRH Learn (http://www.fda.gov/Training/CDRHLearn). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (http://www.fda.gov/DICE) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Mary S. Runner -S

for Tina Kiang, Ph.D.
Acting Director
Division of Anesthesiology,
General Hospital, Respiratory,
Infection Control, and Dental Devices
Office of Device Evaluation
Center for Devices and Radiological Health

Enclosure
Indications for Use

510(k) Number (if known)
K172160

Device Name
Southern Implants PEEK Abutments

Indications for Use (Describe)
The Southern Implants PEEK Abutments are premanufactured prosthetic components directly connected to endosseous dental implants implants and are intended for provisional use up to 180 days as an aid in prosthetic rehabilitation.

Type of Use (Select one or both, as applicable)

☑ Prescription Use (Part 21 CFR 801 Subpart D)  ☐ Over-The-Counter Use (21 CFR 801 Subpart C)

CONTINUE ON A SEPARATE PAGE IF NEEDED.

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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Department of Health and Human Services
Food and Drug Administration
Office of Chief Information Officer
Paperwork Reduction Act (PRA) Staff
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510(k) Summary

Sponsor:
Company Name: Southern Implants (Pty) Ltd.
Company Address: 1 Albert Road
          Irene, South Africa 0062
Telephone: + 27 12 667 1046
Fax: + 27 72 313 5715
Contact Person: Lauranda G. Breytenbach

Summary Preparation Date: February 8, 2018

Device Name:
Trade Name: Southern Implants PEEK Abutments
Common/Usual Name: PEEK Abutments
Classification Name: Abutment, Implant, Dental, Endosseous
Regulation Number: 21 CFR 872.3630
Product Codes: NHA
Device Class: Class II

Predicate Devices:

<table>
<thead>
<tr>
<th>Predicate #</th>
<th>Manufacturer</th>
<th>Device Name</th>
<th>K Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary Predicate</td>
<td>Nobel Biocare AB</td>
<td>NobelProcera PEEK Abutments</td>
<td>K120954</td>
</tr>
</tbody>
</table>
Device Description:

The Southern Implants PEEK abutments are endosseous dental implant abutments. The PEEK abutments are available in a variety of preformed and customized forms. The PEEK abutments are available as healing abutments or temporary abutments.

The healing abutments are attached onto the implant during phase 2 surgery to maintain the tissue opening while the prosthesis is being fabricated. Temporary abutments are used as an aid in manufacturing a temporary prosthesis for prosthetic rehabilitation. They can either be used for direct connection to an endosseous implant or they can be used for connecting the prosthesis to a compact conical abutments. Temporary abutments are available in a 2 mm collar direct to implant and 1mm collar on abutment level. The temporary abutments for the compact conical abutments are for multi-unit use only.

These abutments are made from white PEEK (ASTM F2026).

The PEEK abutments are premanufactured and are available in a variety of connections, engaging and non-engaging, to suit the implant systems manufactured by Southern Implants.

Titanium retaining screws are also available to use with the PEEK abutments.

Indications for Use

The Southern Implants PEEK Abutments are premanufactured prosthetic components directly connected to endosseous dental implants and are intended for provisional use up to 180 days as an aid in prosthetic rehabilitation.

Equivalence to Predicates

Southern Implants (Pty) Ltd submits the information in this Premarket Notification to demonstrate that, for the purposes of FDA’s regulation of medical devices, the subject device is substantially equivalent in indications and design principles to the following legally marketed predicate devices:

K120954, Nobel Procera PEEK Abutments

Tables comparing the subject device and primary predicate devices is provided below.

<table>
<thead>
<tr>
<th>Device and Manufacturer</th>
<th>New device</th>
<th>Primary Predicate (K120954)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Southern Implants PEEK abutments</strong></td>
<td>PK-MAX7-9; PK-MAX7-11; PK-MAX8-11; PK-L-MAX7-9; PK-L-MAX7-11; PK-L-MAX8-9; PK-L-MAX8-11; PKA-Z6-9; PKA28-11</td>
<td>Nobel Procera PEEK Abutments</td>
</tr>
<tr>
<td><strong>Product Code</strong></td>
<td>37819; 37820</td>
<td></td>
</tr>
</tbody>
</table>
Indications for Use
The Southern Implants PEEK Abutments are premanufactured prosthetic components directly connected to endosseous dental implants and are intended for provisional use up to 180 days as an aid in prosthetic rehabilitation.

The Nobel Biocare PEEK Abutments are premanufactured prosthetic components directly connected to endosseous dental implants and are intended for provisional use up to 180 days as an aid in prosthetic rehabilitation.

Intended Use
Healing Abutment

Material
Polyetheretherketone (PEEK)

Colour
White

Attachment Method
Screw retained

Duration of use
180 days, single use

Implant/Abutment Connection
Southern Implants MAX
• External Hex
• Tri-Nex
• Internal Hex

Nobel Biocare
• Internal Conical

Height (mm)
5 – 6 mm

Diameter (mm)
6 x 9
8 x 11

Abutment Profile
Various anatomical shapes intended for different tooth locations

Sterility
Sterile

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Device and Manufacturer

<table>
<thead>
<tr>
<th>New device</th>
<th>Primary Predicate (K120954)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Southern Implants PEEK abutments</td>
<td>NobelProcura PEEK Abutments</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Product Code</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>PKPN2H; PBKBN2NH; PKB2H; PKBNH; PKBA2NH; PKBBB2NH; PKMAX2-2H; PKC-EL-35-2; PKC-EL-43-2; PKC-EL-50-2; PKC-EL-60-2; PKC-DC3-2; PKC-DC4-2; PKC-DCE2; PKC-NDC1-2; ITS6-PKC2; ITS-PKC2; PKC-MC; PKC-MCW.</td>
<td>30259, 30260, 30261, 30262, 30256, 30257, 30258, 31350, 31351, 31352, 31353, 1048-0, 1050-0, 1052-0.</td>
</tr>
</tbody>
</table>

Indications for Use
The Southern Implants PEEK Abutments are premanufactured prosthetic components directly connected to endosseous dental implants and are intended for provisional use up to 180 days as an aid in prosthetic rehabilitation.

The Nobel Biocare PEEK Abutments are premanufactured prosthetic components directly connected to endosseous dental implants and are intended for provisional use up to 180 days as an aid in prosthetic rehabilitation.

Intended Use
Temporary Abutment

Provisional Abutment

Material
Polyetheretherketone (PEEK)
### Southern Implants PEEK Abutments

#### K172160

<table>
<thead>
<tr>
<th>Colour</th>
<th>White</th>
<th>Natural and White</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Attachment Method</strong></td>
<td>Screw retained</td>
<td>Screw retained</td>
</tr>
<tr>
<td></td>
<td>Engaging and Non-Engaging</td>
<td>Engaging and Non-Engaging</td>
</tr>
<tr>
<td><strong>Duration of use</strong></td>
<td>180 days, single use</td>
<td>180 days, single use</td>
</tr>
<tr>
<td><strong>Implant/Abutment</strong></td>
<td>Southern Implants MAX</td>
<td>Nobel Biocare</td>
</tr>
<tr>
<td><strong>Connection</strong></td>
<td>• External Hex</td>
<td>• External Hex</td>
</tr>
<tr>
<td></td>
<td>• Tri-Nex</td>
<td>• Internal Tri-lobe</td>
</tr>
<tr>
<td></td>
<td>• Deep Conical</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Internal Hex</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Internal Taper</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Compact Conical</td>
<td></td>
</tr>
<tr>
<td><strong>Height (mm)</strong></td>
<td>9-14.5</td>
<td>12</td>
</tr>
<tr>
<td><strong>Collar height</strong></td>
<td>1-2</td>
<td>1-1.5</td>
</tr>
<tr>
<td><strong>Diameter (mm)</strong></td>
<td>3.35 – 7.35</td>
<td>Not given</td>
</tr>
<tr>
<td><strong>Abutment Profile</strong></td>
<td>Cylinder with retention feature</td>
<td>Cylinder with retention feature</td>
</tr>
<tr>
<td><strong>Sterility</strong></td>
<td>Sterile</td>
<td>Non-Sterile</td>
</tr>
</tbody>
</table>

The Southern Implants PEEK Healing Abutments minimum diameter and height is the same as the Primary Predicate. The Southern Implants range has a slightly higher version and wider version because it is used on wider implant platforms (up to Ø7 compared to the Ø5.5 platform of the predicate).

The Southern Implants PEEK Temporary Abutments have a range of lengths that falls on either side of the Primary Predicate temporary abutment lengths, by up to 3 mm. The difference in length is immaterial since all lengths fall within the range of lengths that a prosthetic can be made to. If the PEEK abutment is shorter it will just need to be built up more, and if it is longer it will just need to be cut down more.

The Southern Implants PEEK abutments are substantially equivalent to predicate device in:

- Indications for Use
- Intended Use
- Material
- Colour
- Attachment Method
- Duration of Use
- Abutment Profile

The Southern Implants PEEK Abutments are delivered to the patients sterile and is not intended for user sterilization as the predicate. The Southern Implants PEEK Abutments are also available in more connection systems as the predicate to compliment the Southern Implants’ range.

### Nonclinical Testing and Performance Testing

Non-clinical testing data provided or referenced to demonstrate substantial equivalence included:

- The sterilization method is Gamma radiation and has been validated in accordance with ISO 11137.
- The proposed user sterilization method is steam sterilization and is validated in accordance with ISO 17665.
- Packaging was validated in accordance with ISO 11607. Accelerated aging per ASTM-F-1980.
has been applied on the final packaging followed by validating durability to peel and dye tests conditions, in order to substantiate 5 years shelf life.

- The subject device is manufactured from the same material using the same manufacturing method as the predicate, has the same intended use, and the same patient contact type and duration. The subject devices are biocompatible in accordance with ISO 10993-1.
- The subject device shown no cytotoxicity when tested in accordance with ISO 10993-5.

**Clinical Studies**

No clinical studies were conducted.

**Final Conclusion**

Substantial equivalence has been shown for Southern Implants PEEK Abutments.